

Engines of the brain: The computational instruction set of human cognition

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Abstract

Cognition is the action and interaction of multiple brain regions, and these are becoming understood computationally. Simulation and analysis has led to derivation of a set of elemental operations that emerge from individual and combined brain circuits, such that each circuit contributes a particular algorithm, and pairs and larger groups interact to compose further algorithms. We forward the hypothesis that these operations constitute the “instruction set” of the brain, i.e., the basic mental operations from which all other behavioral and cognitive operations are constructed, establishing a unified formalism for description of specialized operations ranging from perception and learning to reasoning and language.

Introduction

“Computation” has come to be associated with computers, but the field’s founders recognized throughout that what they were actually characterizing was the nature of thought. There is still no formal specification for human (or other animal) intelligence; we pursue it solely by imitation of the existence proof, ourselves. The fields of neural networks and artificial intelligence, despite their different tools, usually take as their starting point some behavior of an organism, with the goal of identifying candidate mechanisms that might underlie the described behavior. We present here the converse: by studying the structural (anatomical) and functional (physiological) mechanisms of particular brain structures, the operations that emerge from them may be identified via bottom-up analysis, without pre-specification. The resulting algorithms arising from brain circuit analysis often have unforeseen characteristics, including hierarchical structure, embedded sequences, and even hash coding (see, e.g., Granger et al., 1994; Kilborn et al., 1996; Shimono et al., 2000; Rodriguez et al., 2004). The anatomical layout of these circuits in turn establishes how the individual operators are composed into larger routines. It is hypothesized that these operators, comprising the instruction set of the brain, constitute the basic mental procedures from which all behavioral and cognitive operations are assembled. The resulting constructs give rise to unexpected, and unexpectedly powerful, approaches to complex problems ranging from perception to higher cognition.

Telencephalic Organization

Figure 1 depicts the primary elements of the mammalian forebrain (telencephalon), shared across all mammalian species. Whereas posterior cortex (PC) receives sensory inputs (via T, dorsal thalamus), anterior cortex (AC) produces motor outputs and, in so doing, interacts closely with the elements of the striatal complex (S, striatum; and P pallidum), a more ancient structure that dominates reptilian brains. Both anterior and posterior cortex interact with limbic (hippocampal and amygdaloid) structures (not shown). Mammalian brains scale across several orders of magnitude (from milligrams to kilograms), yet overwhelmingly retain their structural design characteristics. As the ratio of brain size to body size grows, particular allometric changes occur, defining differences between bigger and smaller brain designs. Figure 1b illustrates the three largest changes:

- 1) Connection pathways between anterior and posterior cortex (“fasciculi”) grow large
- 2) Output pathways from striatal complex change relative size: the recurrent pathway back to cortex via thalamus increases relative to the descending motor pathway
- 3) Descending output from anterior cortex to brainstem motor systems (pyramidal tract) grows large

These changes grow disproportionately with increased brain-body ratio, becoming notably outsized in humans. In relatively small-brained mammals such as mice, the primary motor area of neocortex is an adjunct to the striatally driven motor system. Whereas damage to motor cortex in mice causes subtle behavioral motor impairments, damage to motor cortex in humans causes paralysis. In this example of encephalization of function (Jackson, 1925; Ferrier, 1876; Karten, 1991, Aboitiz, 1993) motor operations are increasingly ‘taken over’ by cortex as the size of the pyramidal tract overtakes that of the descending striatal system. In mammals with large brain-body ratios, the role of the striatal complex is presumably altered to reflect that its primary inputs and outputs are now anterior neocortex; in other words, it is now primarily a tool or “subroutine” available for query by anterior cortex. Its operations then are most profitably analyzed in light of its dual utility as organizer of complex motor sequences (in small brained mammals) and as informant to anterior cortex (in large brained mammals).

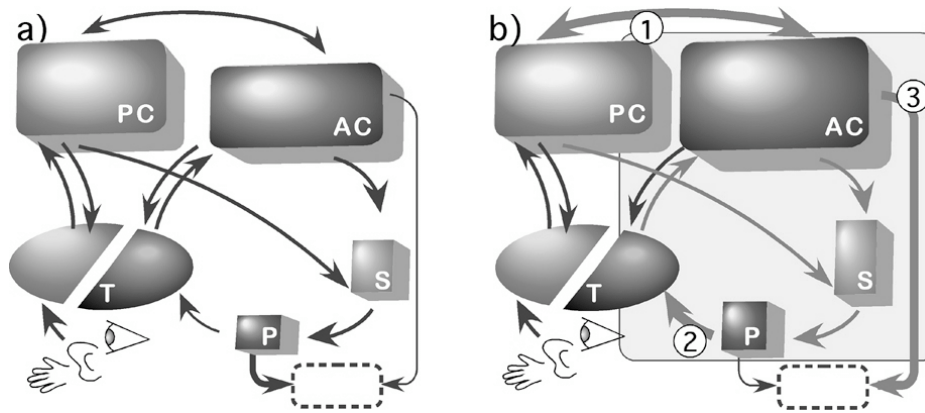


Figure 1: Telencephalic organization

Striatal Complex

The striatal complex or basal ganglia, the primary brain system in reptiles, is a collection of disparate but interacting structures. Figure 2 schematically illustrates the primary components included in the modeling efforts described herein. Distinct components of the basal ganglia exhibit different, apparently specialized designs: matrisomes (matrix), striosomes (patch; which exist as small ‘islands’ embedded throughout the surrounding

matrix regions), globus pallidus, pars interna and externa (GPe and GPi; or pallidum), subthalamic nucleus (STN), tonically active cholinergic neurons (TANs), and substantia nigra pars compacta (SNc). These are connected via a set of varied neurotransmitter pathways including GABA, glutamate (Glu), dopamine (DA), acetylcholine (ACh), and Substance P (Sp) among others, each affecting multiple receptor subtypes.

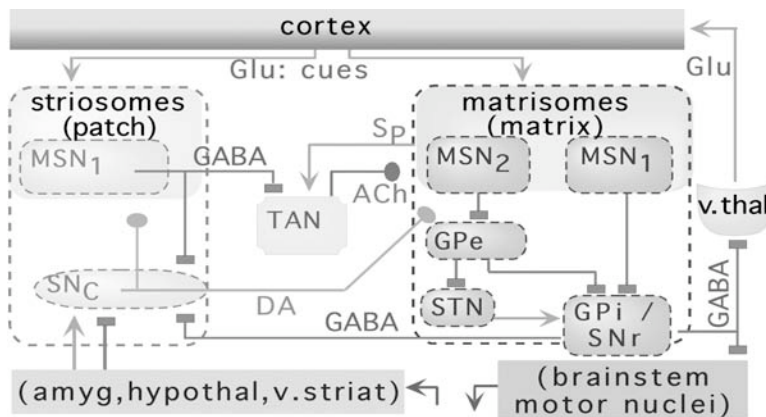


Figure 2. Basal ganglia / striatal complex

The two pathways from cortex through the matrix components of the striatal complex involve different subpopulations of cells in matrisomes: i) MSN1 neurons project to globus pallidus pars interna (GPi), which in turn project to ventral thalamus and back to cortex; ii) MSN2 neurons project to globus pallidus pars externa (GPe), which in turn projects to GPi (and thence to thalamus and cortex). MSN and GP projections are GABAergic, inhibiting their targets. Thus cortical glutamatergic activation of MSN1 cells causes inhibition of GPi cells, which otherwise inhibit thalamic and brainstem targets; hence MSN1 cell activation disinhibits, or enhances, cortical and brainstem activity. In contrast, an extra GABAergic link is intercalated in the pathway from MSN2 neurons to the output stages of matrix; thus

activation of MSN2 neurons decreases activation of cortex and of brainstem nuclei. The pathways from MSN1 and MSN2 neurons are thus termed “go” and “stop,” respectively, for their opposing effects on their ultimate motor and cortical targets. Coordinated operation over time of these pathways can yield a complex combination of activated (go) and withheld (stop) motor responses (e.g., to stand, walk, throw), or correspondingly complex “thought” (cortical) responses.

Two primary afferents to striosomes are cortical and ascending inputs. The former are the same as the inputs to matrix (despite the schematized depiction in the figure, patch components are distributed through, and colocalized with, matrix). Ascending inputs to patch denote “reward” and “punishment” information and have been shown to

up- and down-regulate dopamine from SNc (as well as other dopaminergic sites) in response to external stimuli carrying innate or learned valences (e.g., water to a thirsty organism). A cortically triggered action, followed by an ascending DA reward signal from SNc to patch, selectively enhances active cortical glutamatergic synapses on both matrix and patch targets. Patch output back to SNc then inhibits DA response, so that increased cortical activation of patch (via enhanced synaptic contacts) will come to limit the DA input from SNc. On any given trial, then, the size of the DA signal from SNc reflects the size of the actual ascending DA input (i.e., the reward signal) that occurred over previous trials. Thus with repeated experience, adaptive changes occur in both matrix and patch: initially-random matrix responses to a cortical input become increasingly selected for responses that produce reward, and initial naïve striosomal responses will become increasingly good “predictors” of the size of the reward expected to ensue as a result of the action.

Tonically active cholinergic neurons (TANs) represent a small fraction (< 5%) of the number of cells in the striatal complex yet densely contact cells throughout matrix; thus they likely play a modulatory role rather than

conveying specific information. The GABAergic inhibition of these cells by patch will come to increase for those patch responses that lead to reward, since in these instances the cortical drivers of these patch responses become synaptically enhanced. Thus in those circumstances where cortical inputs lead to expected reward, TANs will tend to have less excitatory effect on matrix. Since the TAN afferents to matrix are dense and nontopographic, they represent a random “background noise” input, which can increase variance in selected matrix responses to cortical inputs, making the striatally-selected motor response to a cortical input somewhat nondeterministic. The resulting behavior should appear “exploratory,” involving a range of different responses to a given stimulus. Synaptic increases in patch, in addition to causing accurate “predictions” of reward size, as described, also increasingly inhibit TANs, diminishing the breadth of exploratory variability. Thus as rewards occur, not only will reward-associated responses be increasingly selected by matrix, but the variability among those responses will decrease. Analyses suggest detailed comparisons of basal ganglia and standard reinforcement learning systems (Schultz et al., 1997; Dayan et al., 2000; see Table 1).

Table 1. Simplified basal ganglia algorithm

- 1) Choose action A. Set reward_estimate \leftarrow 0
Set max_randomness \leftarrow R > 0
- 2) randomness \leftarrow max_randomness – reward_estimate
- 3) reward \leftarrow Eval(A + randomness)
- 4) If reward > reward_estimate then
A \leftarrow A + randomness
reward_estimate \leftarrow reward
- 5) goto step 2)

Neocortex / Thalamocortical system

Neurons throughout neocortex are organized into relatively stereotypical architectures (Figure 3a). Although cortical studies describe some (subtle but potentially crucial) differences among various cortical regions (Galuske et al., 2000; Gazzaniga, 2000), the overwhelmingly shared characteristics justify attempts to identify common basic functionality, which may be augmented by special purpose capabilities in some regions (Lorente de No, 1938; Szentagothai, 1975; Keller & White, 1989; Rockel et al., 1980; Castro-Alamancos & Connors, 1997; Braitenberg & Schuz, 1998; Valverde, 2002).

Two parallel circuit types occur, involving topographic projections of certain restricted thalamic populations and broad, diffuse projections from the remaining thalamic neurons. These two populations of thalamic cells, respectively termed “core” and “matrix” (no relation, confusingly enough, with “matrix” in striatum), are distinguishable by their targets, topography,

and chemistries (Jones, 2001). The topographically organized projections from thalamic core synapse largely on layer IV and deep layer III cells (Fig 3b); the diffuse matrix projections form synapses predominantly in layer I, on the apical dendrites of layer II, III and V cells (Fig 3c). (Although the topographic afferents to middle cortical layers, e.g. LGN to visual cortex, are often thought of as the primary input to sensory neocortex, these fibers actually comprise only about 6% of the synapses onto their primary layer IV targets, with the remaining afferents coming largely from lateral cortico-cortical connections (Freund et al., 1985; 1989; Peters & Payne, 1993).

Peripheral inputs activate thalamic core cells, which in turn participate in topographic activation of middle cortical layers; e.g., ear \rightarrow cochlea \rightarrow auditory brainstem nuclei \rightarrow ventral subdivision of medial geniculate (MGv) \rightarrow primary auditory cortex (A1). In contrast, matrix nuclei are most strongly driven by corticothalamic feedback (Diamond et al. 1992). Thus peripheral afferents

activate core nuclei, which activate cortex (via a stereotypical vertically organized pattern: middle layers → superficial → deep layers), then activating core and matrix nuclei via corticothalamic projections (Mountcastle 1957; Hubel & Wiesel 1977; Di et al. 1990).

Three primary modes of activity have typically been reported for thalamic neurons: tonic, rhythmic and arrhythmic bursting. The latter appears predominantly during non-REM sleep whereas the first two appear during waking behavior (McCarley et al., 1983; Steriade & Llinas, 1988; McCormick & Bal, 1994). There is strong evidence for ascending influences (e.g., basal forebrain) affecting the probability of neuronal response during the peaks and troughs of such “clocked” cycles. The most excitable cells will tend to fire in response even to slight afferent activity whereas less excitable neurons will only be added in response to stronger input; this excitability gradient selectively determines the order in which neurons will be recruited to respond to inputs of any given intensity.

Axons of inhibitory interneurons densely terminate preferentially on the bodies, initial axon segments, and proximal apical dendrites of excitatory pyramidal cells in cortex, and thus are well situated to exert powerful control over the activity of target excitatory neurons. When a field of excitatory neurons receives afferent stimulation, those that are most responsive will activate the local inhibitory cells in their neighborhood, which will in turn inhibit local excitatory cells. The typical time course of an excitatory (depolarizing) postsynaptic potential (PSP) at normal resting potential, in vivo, is brief (15-20 msec), whereas corresponding GABAergic inhibitory PSPs can last roughly an order of magnitude longer (80-150 msec) (Castro-Alamancos and Connors, 1997). Thus excitation tends to be brief, sparse, and curtailed by longer and stronger feedback lateral inhibition (Coultrip et al., 1992).

Based on the biological regularities specified, a greatly simplified set of operations has been posited (Rodriguez et al., 2004). Distinct algorithms arise from simulation and analysis of core vs. matrix loops (see Tables 2 & 3).

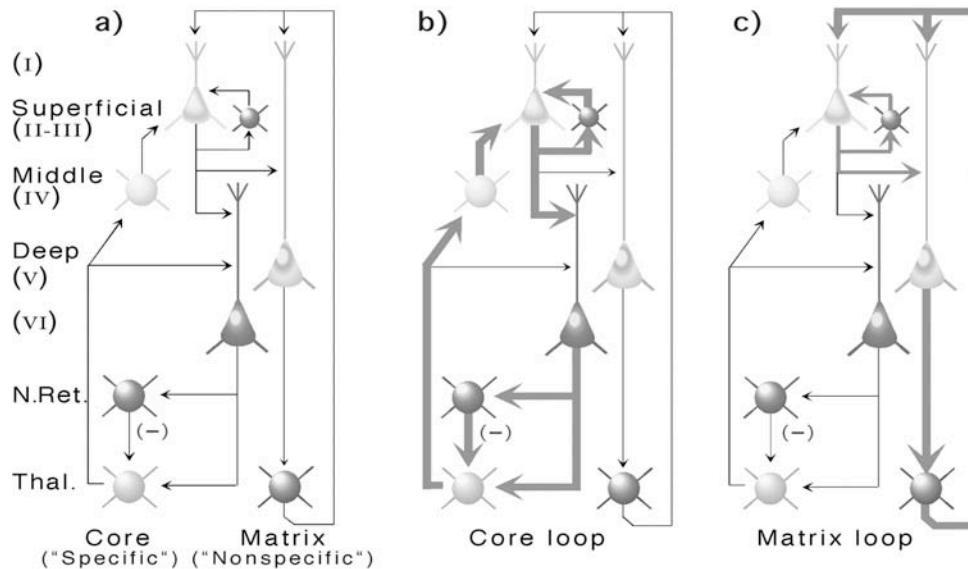


Figure 3. Thalamocortical loops

Thalamocortical “core” circuits. In the core loop, simulated superficial cells that initially respond to a particular input pattern become increasingly responsive not only to that input but also to a range of similar inputs (those that share many active lines; i.e., small Hamming distances from each other), such that similar but distinguishable inputs will come to elicit identical patterns of layer II-III cell output, even though these inputs would have given rise to slightly different output patterns before synaptic potentiation. These effects can be described in terms of the formal operation of clustering, in which sufficiently similar inputs are placed into a single category or cluster. This can yield useful generalization

properties, but somewhat counterintuitively, it prevents the system from making fine distinctions among members of a cluster. For instance, four similar inputs may initially elicit four slightly different patterns of cell firing activity in layer II-III cells but after repeated learning / synaptic potentiation episodes, all four inputs may elicit identical cortical activation patterns. Results of this kind have been obtained in a number of different models with related characteristics (von der Malsburg, 1973; Grossberg, 1976; Rumelhart & Zipser, 1985; Coultrip et al., 1992).

Superficial layer responses activate deep layers. Output from layer VI initiates feedback activation of nucleus reticularis (N.Ret) (Liu and Jones 1999), which in

turn inhibits the core thalamic nucleus (Fig 3b). Since, as described, topography is preserved through this sequence of projections, the portions of the core nucleus that become inhibited will correspond topographically to those portions of L.II-III that were active. On the next cycle of thalamocortical activity, the input will arrive at the core against the background of the inhibitory feedback from N.Ret, which has been shown to last for hundreds of milliseconds (Cox et al., 1997; Zhang et al., 1997). Thus it is hypothesized that the predominant component of the next input to cortex is only the un-inhibited remainder of the input, whereupon the same operations as before are performed. Thus the second cortical response will consist of a quite distinct set of neurons from the initial response, since many of the input components giving rise to that first response are now inhibited. Analysis of the second (and ensuing) responses in computational models has shown successive sub-clustering of an input: the first cycle of response identifies the input's membership in a general category of similar objects (e.g., flowers), the next response (a fraction of a second later) identifies its membership in a particular subcluster (e.g., thin flowers; flowers missing a petal), then sub-sub-cluster, etc. Thus the system repetitively samples across time, differentially activating specific target neurons at successive time points, to discriminate among inputs. An initial version of this derived algorithm arose from studies of feedforward excitation and feedback inhibition in the olfactory paleocortex and bulb, and was readily generalized to non-olfactory modalities (vision, audition) whose superficial layers are closely related to those of

olfactory cortex, evolutionarily and structurally. The method can be characterized as an algorithm (Table 2).

Analysis reveals the algorithm's time and space costs. The three time costs for processing of a given input X are: i) summation of inputs on dendrites; ii) computation of "winning" (responding) cells C ; iii) synaptic weight modification. For n learned inputs of dimensionality N , in a serial processor, summation is performed in $O(nN)$ time, computation of winners takes $O(n)$ time, and weight modification is $O(N \log n)$. With appropriate parallel hardware, these three times reduce to $O(\log N)$, $O(\log n)$, and constant time respectively, i.e., better than linear time. Space costs are similarly calculated: given a weight matrix W , to achieve complete separability of n cues, the bottom of the constructed hierarchy will contain at least n units, as the leaves of a tree with $\log Bn$ hierarchical layers, where B is the average branching factor at each level. Thus the complete hierarchy will contain $\sim n[B/(B-1)]$ units, i.e., requiring linear space to learn n cues (Rodriguez et al., 2004).

These costs compare favorably with those in the (extensive) literature on such methods (Ambros-Ingerson et al., 1990; Gluck & Granger, 1993; Kilborn et al., 1996; Rodriguez et al., 2004). Elaboration of the algorithm has given rise to families of computational signal processing methods whose performance on complex signal classification tasks has consistently equaled or outperformed those of comparable methods (Coultrip and Granger, 1994; Kowtha et al., 1994; Granger et al., 1997; Benvenuto et al., 2002; Rodriguez et al., 2004).

Table 2. Thalamocortical Core Algorithm

```
for input X
  for C ∈ win(X, W)
    Wj ← Wj + k(X - C)
  end_for
X ← X - mean(win(X, W))
end_for
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where

X = input activity pattern (vector);

W = layer I synaptic weight matrix;

C = responding superficial layer cells (col vector);

k = learning rate parameter;

$\text{win}(X, W)$ = column vector in W most responsive to X before lateral inhibition [$\forall j, \max(X \cdot W_j)$]

Thalamocortical "matrix" circuits. In contrast to the topography-preserving projections in the "core" loop between core thalamus and cortex, the diffuse projections from layer V to matrix nuclei, and from matrix nuclei back to cortex (Fig 3c) are modeled as sparsifying and orthogonalizing their inputs, such that any structural relationships that may obtain among inputs are not retained in the resulting projections. Thus input patterns in matrix or in layer V that are similar may result in very

different output patterns, and vice versa. As has been shown in previously published studies, due to the nontopographic nature of layer V and matrix thalamus, synapses in layer V are very sparsely selected to potentiate, i.e., relatively few storage locations (synapses) are used per storage/learning event (Granger et al., 1994; Aleksandrovsky et al., 1996; Rodriguez et al., 2004). For purposes of analysis, synapses are assumed to be binary (i.e., assume the lowest possible precision: synapses that

are either naïve or potentiated). A sequence of length L elicits a pattern of response according to the algorithm given previously for superficial layer cells. Each activated superficial cell C in turn activates deep layer cells. Feedforward activity from the matrix thalamic nucleus also activates layer V . Synapses on cells activated by both sources (the intersection of the two inputs) become potentiated, and the activity pattern in layer V is fed back to matrix. The loop repeats for each of the L items in the sequence, with the input activity from each item interacting with the activity in matrix from the previous step (see Rodriguez et al., 2004).

Activation of layer V in rapid sequence via superficial layers (in response to an element of a sequence) and via matrix thalamus (corresponding to feedback from a previous element in a sequence) selects responding cells sparsely from the most activated cells in the layer (Coultrip et al., 1992) and selects synapses on those cells sparsely as a function of the sequential pattern of inputs arriving at the cells. Thus the synapses potentiated at a given step in layer V correspond both to the input occurring at that time step together with orthogonalized feedback arising from the input just prior to that time step. The overall effect is “chaining” of elements in the input sequence via the “links” created due to coincident layer V activity corresponding to current and prior input elements. The sparse synaptic potentiation enables layer V cells to act as a novelty detector, selectively responding to those sequential strings that have previously been presented (Granger et al., 1994). The implicit data structures created are trees in which initial sequence elements branch to their multiple possible

continuations (“tries,” Knuth, 1997). Sufficient information therefore exists in the stored memories to permit completion of arbitrarily long sequences from just prefixes that uniquely identify the sequence. Thus the sequence “Once upon a time” may elicit (or “prime”) many possible continuations whereas “Four score and seven” elicits a specific continuation.

The resulting algorithm (see Table 3) can be characterized in terms of computational storage methods that are used when the number of actual items that occur are far fewer than those that in principle could occur. The number of possible eight-letter sequences in English is $26^8 \approx 200,000,000,000$, yet the eight-letter words that actually occur in English number less than 10,000, i.e., fewer than one ten-millionth of the possible words. The method belongs to the family of widely-used and well-studied data storage techniques of “scatter storage” or “hash” functions, known for the ability to store large amounts of data with great efficiency. Both analytical results and empirical studies have found that the derived matrix loop method requires an average of less than two bits (e.g., just two low-precision synapses) per complex item of information stored. The method exhibits storage and successful retrieval of very large amounts of information at this rate of storage requirement, leading to extremely high estimates of the storage capacity of even small regions of cortex. Moreover, the space complexity of the algorithm is linear, or $O(nN)$ for n input strings of dimensionality N ; i.e., the required storage grows linearly with the number of strings to be stored (Granger et al., 1994; Aleksandrovsky et al., 1996; Rodriguez et al., 2004).

Table 3. Thalamocortical Matrix Algorithm

```

for input sequence X(L)
  for C ∈ TopographicSuperficialResponse(X(L))
    for V(s) ∈ C ∩ NNtResponse(X(L-1))
      Potentiate( V(s) )
    NNt(L) ← NontopographicDeepResponse(V)
  end_for
end_for
end_for

```

where L = length of input sequence;

C = columnar modules activated at step $X(L)$;

$V(s)$ = synaptic vector of responding layer V cell,

$NNt(L)$ = response of nonspecific thalamic nucleus to feedback from layer V .

Combined telencephalic algorithm operation and the emergence of complex specializations. In combination with time dilation and compression algorithms arising from amygdala and hippocampal models (Granger & Lynch, 1991; Granger et al., 1994; Kilborn et al., 1996), a rich range of operations is available for composition into complex behaviors. From the operation of thalamocortical loops arises the learning of similarity-

based clusters (Table 2) and brief sequences (Table 3), yielding the primary data structure of thalamocortical circuitry: sequences of clusters. These are embedded into thalamo-cortico-striatal (TCS) loops which enable reinforcement-based learning of these sequences of clusters. The output of any given cortical area becomes input (divergent and convergent) to other cortical areas, as well as receiving feedback from those cortical areas.

Each such region in the TCS architecture performs the same processing on its inputs, generating learned nested sequences of clusters of sequences of clusters.

Auditory cue processing. Figure 4a illustrates a spectrogram (simplified cochleogram) of a voice stream (the spoken word “blue”), as might be processed by

presumed auditory “front end” input structures. Proceeding left to right (i.e., in temporal order) and identifying “edges” that are readily detected (by simple thresholding) leads to creation of brief sequences / segments corresponding to these edges as in Figure 4b.

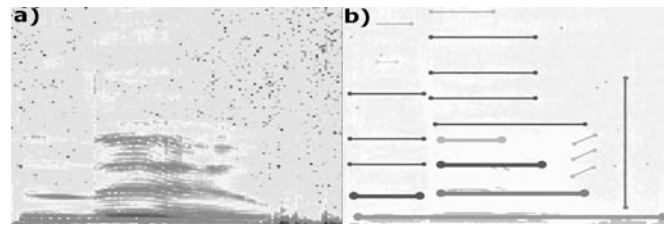


Figure 4. Spectrogram and learned cortical sequences

The learned cortical sequences (characterized as line segments) correspond to constituents of the signal. As multiple instances of the signal are learned, some features will be strengthened more than others, corresponding to a statistical average of the signal rather than of any specific instance. Outputs from cortical areas are input to other cortical areas, combining individual pairwise sequences into sequences of sequences (actually sequences of clusters of sequences of clusters, etc.), and statistics are accreted for these by the same mechanisms. The result is a widely distributed set of synaptic weights that arise as a result of training on instances of this kind. (There is contention in the literature as to whether such learned internal patterns of synaptic weights are “representations,” a term that has baggage from other fields. Without engaging this controversy, we use the expression as a term of convenience for these patterns of weights.) These differ from many other types of representations, in that they are not strict images of their inputs but rather are statistical “filters” that note their sequence of features (or sequence of sequences) in a novel input, and compete against other feature filters to identify a “best partial match” to the input. It is notable that since each sequence pair simply defines relative

positions between the pair, they are independent of particular frequencies or exact time durations.

Figure 5 illustrates two different instances of the utterance “blue” that, after learning, can be recognized by the algorithm as members of the same category, since they contain many of the same organization of grammatical elements (sequences of clusters, and sequences of clusters of sequences of clusters), whereas other utterances contain distinguishing differences. These representations, arising simply from distributed patterns of synaptic strengthening in the described brain circuit networks, have desirable properties for recognition tasks.

The “best partial match” process can pick out candidate matches from a stream of inputs. Thus the detector for “blue” and that for “bird” identify their respective targets in a continuous utterance (e.g., “the blue bird”). Recognition systems traditionally have difficulty with segmentation, i.e., division of a stream into parts. In the proposed recognition scheme, recognition and segmentation are iteratively interleaved: identification of the sequence components of a candidate word in the stream gives rise to a candidate segmentation of the stream. Competing segmentations (e.g., from sequence components of other words overlapping) may overrule one segmentation in favor of an alternative.

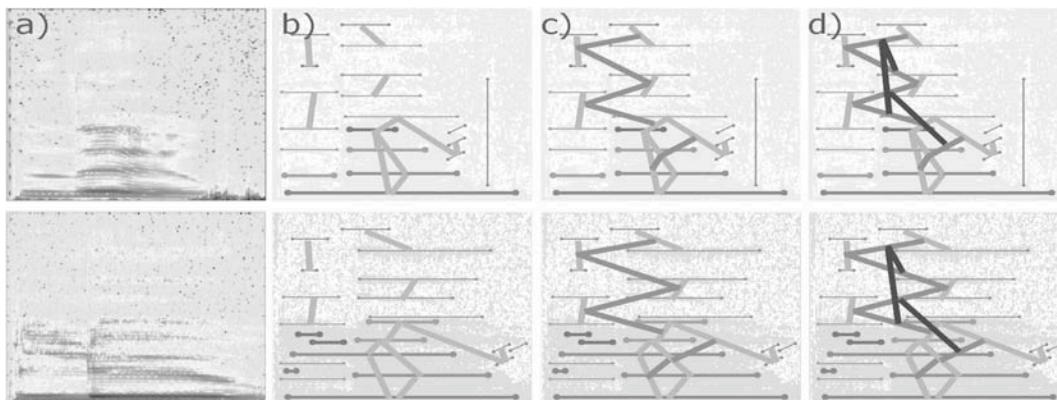


Figure 5. Two utterances and illustration of learned nested sequences

The figure illustrates the nested nature of the operation of the thalamo-cortico-striatal loops. Initial processing of input a) involves special-purpose “front ends” that in the model are carried out by (well-studied) Gabor filters and edge detection methods, producing a first internal representation of sequences as seen in Figure 4. Each successive stage of processing takes as input some combination of the outputs of prior stages. Thus the brief sequences in Figure 4b become input to a copy of the same mechanism, which identifies sequences of the sequences (5b). Downstream regions then identify sequences of those sequences, and so on (5c,d). With learning, the resulting set of relative feature positions comes to share substantial commonalities that are partial-matched, as in the two different utterances of the word “blue in the top and bottom frames of Figure 5.

Visual image processing. Once past the initial, specialized “primary” cortical sensory regions, thalamocortical circuits are remarkably similar (though, as mentioned, differences have been found, with unknown implications). Moreover, the vast majority of cortical areas appear to receive inputs not originating just from a single sensory modality but from conjunctions of two or more, begging the question of whether different internal “representations” can possibly be used for different modalities. Auditory cortical regions arise relatively early

in mammalian evolution (consistent with the utility of non-visual distance senses for nocturnal animals) and may serve as prototypes for further cortical elaboration, including downstream (non-primary) visual areas. It is here hypothesized that, although primary cortical regions perform specialized processing, subsequent cortical regions treat all inputs the same, regardless of modality of origin. The physiological literature suggests particular visual front end processing (arising from retina, LGN, early cortical areas) resulting in oriented line and curve segments comprising an image. From there on, images may be processed as sounds, though due to recruitment of front end visual processing, arbitrary covert “movements” through an image are assumed to occur, rather than processing being limited to an arbitrary “left to right” corresponding to the flow of time in an auditory image. I.e., it is as though auditory processing were a callable subroutine of visual processing. Thus, after initial processing of an image (such as part of Figure 6a) (performed in this case via oriented Gabor filters (6b) at different frequency parameters, to roughly approximate what has been reported for visual front end processing from many sources over many years), the resulting segments (pairwise sequences) are composed into sequences of sequences (6c), etc until, over training trials, they become statistical representations of the objects (e.g., letters) on which they have been trained (6d).

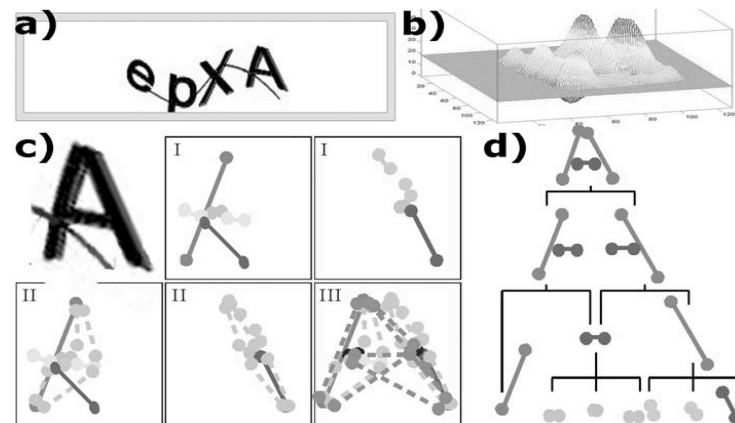


Figure 6. Nested sequences of clusters identified in images (see text).

As with auditory data, this method leads to representations that iteratively alternate recognition and segmentation; i.e., there exists no separate segmentation step but rather candidate segments emerge, as recognizers compete to identify best partial matches in an image. Further characteristics shared with auditory processing include a number of invariances: translation, scaling and distortion, as well as resistance to partial occlusion. Again, these invariances are not add-on processing

routines but rather emerge as a result of the processing. Since the sequences, and sequences of sequences, record relative relationships as opposed to absolute locations, and since the front end filtering occurs across multiple size and frequency scales, recognition of a small A in a corner proceeds just like that of a large centered A. And since the result is merely a best partial match, rather than a complete match, a partially occluded or distorted A may match to within threshold (Figure 7).



Figure 7. Emergent invariances from the derived methods (distortion, scaling, partial occlusion).

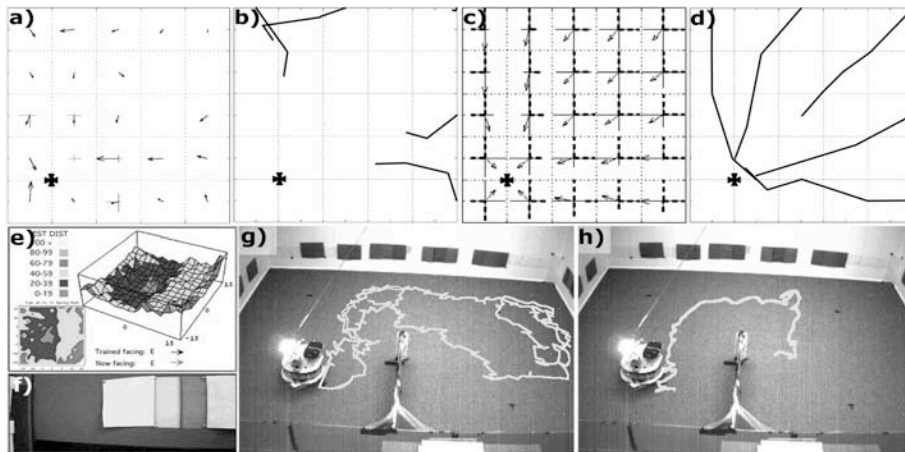


Figure 8. Learned internal representations, and trajectories.

Navigation. Presentation of locations containing a hard-coded artificial desirable “goal” state, and sequential reinforcement training from various starting locations, causes the system to improve its approaches to the goal from arbitrary starting points. Figure 8 shows the internal representations (a,c) constructed in the striatal complex as a result of training trials, and illustrates sample trajectories (b,d) to the goal from five starting points, both before (a,b) and after (c,d) this training. The representations correspond to the learned positive and negative “strengths” of four candidate movement directions (N,S,E,W), along with a resultant vector, at each location in the grid. Figures 8e-f show the corresponding internal representation (e) from photographs (f), enabling a robot to learn efficient navigation to a goal from different starting locations (8g,h).

Hierarchical grammatical structure. It is notable that the emergent data structure of the thalamo-cortico-striatal model, nested sequences of clusters, is a superset of the structures that constitute formal grammars (in particular, graph-grammars): i.e., ordered sequences of elements, such that each element (node) represents either a category (in this case a cluster), or expands to another node (nesting). Among the useful properties of these data structures is their incremental growth: adding new copies of TCS loops (as brain size grows) increases the size of the grammars that can be learned, corresponding to the incremental addition of learned “rules” to the grammar. Thus small-brained mammals may acquire relatively small internal grammars, enabling learning of comparatively simple mental constructs, whereas larger-brained mammals may learn increasingly complex

internal representations. This is specifically seen in the examples above. E.g., back in Figure 5, successive processing of the input, carried out by increasingly “downstream” components of the model, identifies first a simple set of features and relations among those features; then successively more complex nested relations, and relations among relations. In the model, though all “regions” are identical in structure, they receive somewhat different (though overlapping) inputs (e.g., certain visual features; certain combinations of visual and auditory features). After exposure to multiple inputs, regional specializations of function (e.g., human voices vs. other sounds; round objects vs. angular objects) arise due to lateral inhibition among areas, giving rise to regions that are performing the same computational function but become increasingly dedicated to the processing of particular types of inputs.

The incremental nature of the data structure (nested sequences of clusters) enables it to grow simply by adding to the components or “rules” acquired by the grammar, adding to the power of the resulting behavior that the grammar can give rise to. As abstractions (nested sequences of clusters of sequences of clusters) are built up through the incremental grammar, slightly more complicated behaviors arise from its use. As more telencephalic “real estate” is added, longer (nested) sequences of clusters emerge, corresponding to more complex syntax. That is, changing nothing of the structure of thalamocortical loops, only the number of them, can give rise to new function.

The extensible (generative) nature of human language has typically been explained in terms of grammars of this kind: from a given grammar, a

potentially infinite number of outputs (strings in the language) can be produced. Humans uniquely exhibit rapidly-acquired complex grammatical linguistic behavior, prompting the search for uniquely human brain regions that could explain the presence of this faculty in humans and its absence in other primates (see, e.g., Hauser et al., 2002; Fitch and Hauser 2004; O'Donnell et al., 2005; Preuss 1995; 2000; Galuske et al., 2000). The modeling described herein leads to a specific hypothesis: that human language arises in the brain as a function of the number of thalamo-cortico-striatal loops. Rather than a wholly new brain module that differentially processes for language, the evolutionary addition of TCS modules, some of which indeed become dedicated to communication, leads to the incremental acquisition of linguistic abilities. This growth need not be linear; grammars have the property of exhibiting apparently new behaviors due to the addition of just a few rules. There are no living primates with brain sizes between those of apes and humans, a fourfold difference; if human language arises directly from increases to these TCS loops, then intermediate prelinguistic abilities may have been present in early hominids, even though not in extant primates. The conjecture is consistent with a broad range of constraints that are argued to rule out alternative hypotheses (see, e.g., Pinker 1999; Pinker and Jackendoff 2005).

The processing of linguistic input, then, is not a different function from that of other brain processing, but rather is just one special case of the computational faculties present in smaller brains. With an understanding of the specific nature of these computations, it is possible to see how they operate on simpler (e.g., perceptual) inputs as well as complex (linguistic) inputs, differing largely in the depth of processing and thus the size of the grammars that are constructed for these inputs.

Conclusions

Human brains arose via a series of intermediaries and under a range of different conditions, without any set of computational plans or top-down principles. Thus brains and their constituent circuits are not "optimized" for any particular task but represent earlier circuits co-opted to perform new jobs, as well as compromises across multiple tasks that a given circuit may have to participate in under different circumstances. Bottom up analysis of circuits, without targeting any "intended" or "optimized" functions, leads to a set of computational functions that may comprise the "instruction set" of a brain's most basic operations, of which other operations are composed. The overwhelming regularity of cortical structures, and of large loops through cortical and striatal telencephalon, suggests the universality of the resulting composite operations.

The basic algorithms that have been derived include many that are not typically included in proposed

"primitive" sets: sequence completion, hierarchical clustering, retrieval trees, hash coding, compression, time dilation, reinforcement learning. Analysis indicates these algorithms' computational efficiency, showing that they scale well as brain size increases (Rodriguez et al., 2004). Application of these derived primitives gives rise to a set of unusual approaches to well-studied tasks ranging from perception to navigation, and illustrates how the same processes, successively re-applied, enable learning of data structures that account for generative human language capabilities.

Persistent questions of brain organization are addressed. For instance, How can replication of roughly the same (neocortical) circuit structure give rise to differences in kind rather than just in number? Thalamocortical and corticostriatal algorithms must be constituted such that making more of them enables interactions that confer more power to larger assemblies. This property is certainly not universal (e.g., backpropagation costs scale as the square of network size, and do not solve new kinds of problems simply by growing larger).

What relationships, if any, exist between early sensory operations and complex cognitive operations? The specific hypothesis is forwarded here that, beyond initial modality-specific "front end" processing, all telencephalic processing shares the same operations arising from successive thalamo-cortico-striatal loops. Complex "representations" (objects, spaces, grammars, relational dictionaries) are composed from simpler ones; "cognitive" operations on these complex objects are the same as the perceptual operations on simpler representations; and grammatical linguistic ability is constructed directly from iterative application of these same operators.

Procedures that seem easy and natural to humans (and, often, to other animals) such as image recognition, sound recognition, tracking, etc., have been notoriously difficult for artificial systems to perform. Many of these tasks are ill-specified, and the only reason that we believe that they are computable is the existence proof that we perform them. Moreover, many systems that learn statistically have been shown to be inadequate to the task of learning rule-like cognitive abilities (Pinker, 1999). We have here illustrated that data structures of grammatical form arise from models that contain the anatomical architectures and physiological operations of actual brain circuits, demonstrating how this class of circuit architecture can avoid the problems of extant "neural network" models and give rise to computational constructs of a power appropriate to the tasks of human cognition. Ongoing bottom-up analyses of brain circuit operation may continue to provide novel engineering approaches applicable to the seemingly intractable problems of cognition.

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References

- Aboitiz F (1993) Further comments on the evolutionary origin of mammalian brain. *Med Hypoth* 41: 409-418.
- Aleksandrovsky B, Whitson J, Garzotto A, Lynch G, Granger R (1996) An algorithm derived from thalamocortical circuitry stores & retrieves temporal sequences. *IEEE Int Conf Patt Rec*, 550-554.
- Ambros J, Granger R, Lynch G (1990) Simulation of paleocortex performs hierarchical clustering. *Science* 247:1344-1348.
- Benvenuto J, Jin Y, Casale M, Lynch G, Granger R (2002) Identification of diagnostic evoked response potential segments in Alzheimer's Disease. *Exper Neurol* 176:269-276.
- Bourassa J, Deschenes M (1995) Corticothalamic projections from the primary visual cortex in rats: a single fiber study using biocytin as an anterograde tracer. *Neuroscience* 66:253-263.
- Braitenberg V, Schüz A (1998) *Cortex: statistics and geometry of neuronal connectivity*, NY: Springer.
- Castro-Alamancos M, Connors B (1997) Thalamocortical synapses. *Prog Neurobiol* 51:581-606.
- Coultrip R, Granger R (1994) LTP learning rules in sparse networks approximate Bayes classifiers via Parzen's method. *Neural Networks* 7:463-476.
- Coultrip R, Granger R, Lynch G (1992) A cortical model of winner-take-all competition via lateral inhibition. *Neural Networks* 5:47-54.
- Cox C, Huguenard J, Prince D (1997) Nucleus reticularis neurons mediate diverse inhibitory effects in thalamus. *PNAS* 94:8854-59.
- Deschenes M, Veinante P, Zhang ZW (1998) The organization of corticothalamic projections: reciprocity versus parity. *Brain Res Brain Res Rev* 28:286-308.
- Di S, Baumgartner C, Barth D (1990) Laminar analysis of extracellular field potentials in rat barrel cortex. *J Neurophysiol* 63:832-840.
- Diamond M, Armstrong-James M, Ebner F (1992) Somatic sensory responses in the rostral sector of the posterior group (POM) and in the ventral posterior medial nucleus (VPM) of the rat thalamus. *J Comp Neurol* 318:462-476.
- Ferrier D (1876) *Functions of the brain*. London: Smith, Elder.
- Fitch T, Hauser M (2004) Computational constraints on syntactic processing in a nonhuman primate. *Science* 303: 377-380.
- Freund T, Martin K, Whitteridge D (1985) Innervation of cat visual areas 17+18 by physiologically identified X & Y-type thalamic afferents. I. Arborization patterns & quantitative distribution of postsynaptic elements. *J Comp Neur* 242:263-274.
- Freund T, Martin K, Soltesz I, Somogyi P, Whitteridge D (1989) Arborisation pattern and postsynaptic targets of physiologically identified thalamocortical afferents in striate cortex of the macaque monkey. *J Comp Neurol* 289:315-336.
- Galuske RA, Schlote W, Bratzke H, Singer W (2000) Interhemispheric asymmetries of the modular structure in human temporal cortex. *Science* 289:1946-1949.
- Gazzaniga MS (2000) Regional differences in cortical organization. *Science* 289:1887-1888.
- Gluck M, Granger R (1993) Computational models of the neural bases of learning and memory. *Ann Rev Neurosci* 16:667-706.
- Granger R, Lynch G (1991) Higher olfactory processes: perceptual learning and memory. *Curr Opin Neurobiol* 1:209-214.
- Granger R, Whitson J, Larson J, Lynch G (1994) Non-Hebbian properties of long-term potentiation enable high-capacity encoding of temporal sequences. *Proc Nat Acad Sci* 91:10104-10108.
- Granger R, Wiebe S, Taketani M, Ambros-Ingerson J, Lynch G (1997) Distinct memory circuits comprising the hippocampal region. *Hippocampus* 6:567-578.
- Granger R (2002) *Neural Computation: Olfactory cortex as a model for telencephalic processing*. In: *Learning & Memory* (J. Byrne, Ed), MacMillan Reference Books, pp. 445-450
- Grossberg S (1976) Adaptive pattern classification and universal recoding. *Biol Cybern* 23:121-134.
- Hauser M, Chomsky N, Fitch T (2002). The language faculty: What is it, who has it, and how did it evolve? *Science* 298: 1569-1579.
- Hubel D, Wiesel T (1977) Functional architecture of macaque monkey visual cortex. *Proc R Soc Lond B Biol Sci* 198:1-59.
- Jackson JH (1925) *Neurological fragments*. London: Oxford Univ
- Jones E (1998) A new view of specific and nonspecific thalamocortical connections. *Adv Neurol* 77:49-71.
- Karten H (1991) Homology and evolutionary origins of 'neocortex'. *Brain Beh Evol*, 38: 264-272.
- Keller A, White E (1989) Triads: a synaptic network component in cerebral cortex. *Brain Res* 496:105-112.
- Kilborn K, Granger R, Lynch G (1996) Effects of LTP on response selectivity of simulated cortical neurons. *J Cog Neuro* 8:338-353.
- Knuth D (1997) *The art of computer programming*. MA: Addison-Wesley.
- Kowtha V, Satyanara P, Granger R, Stenger D (1994) Learning & classification in a noisy environment by a simulated cortical network. *3rd Ann Comp Neur Sys* 245-50. Boston: Kluwer
- Liu X, Jones E (1999) Predominance of corticothalamic synaptic inputs to thalamic reticular nucleus neurons in the rat. *J Comp Neurol* 414:67-79.
- Lorente de No R (1938) Cerebral cortex: Architecture, intracortical connections, motor projections. In: *Physiology of the nervous system* (Fulton J, ed), pp 291-340. London: Oxford.
- McCarley R, Winkelman J, Duffy F (1983) Human cerebral potentials associated with REM sleep rapid eye movements: links to PGO waves and waking potentials. *Brain Res* 274:359-364.
- McCormick D, Bal T (1994) Sensory gating mechanisms of the thalamus. *Curr Opin Neuro* 4:550-556.
- Mountcastle V (1957) Modality and topographic properties of single neurons of cat's somatic sensory cortex. *J Neurophysiol* 20:408-434.

O'Donnell T, Hauser M, Fitch T (2004). Using mathematical models of language experimentally. *Trends Cog Sci*, 9: 284-289.

Peters A, Payne B (1993) Numerical relationships between geniculocortical afferents and pyramidal cell modules in cat primary visual cortex. *Cerebral Cortex* 3:69-78.

Pinker S (1999). *Words and rules: the ingredients of language*. New York: HarperCollins.

Pinker S, Jackendoff R (2005). The faculty of language: what's special about it? *Cognition* (In press).

Preuss T (1995). Do rats have prefrontal cortex? The Rose-Woolsey-Akert program reconsidered. *J Cog Neurosci*, 7: 1-24.

Preuss T (2000). What's human about the human brain? In: *The New Cognitive Neurosciences*. M.Gazzaniga (Ed.), Cambridge, MA: MIT Press, pp.1219-1234.

Rockel AJ, Hiorns RW, Powell TPS (1980) Basic uniformity in structure of the neocortex. *Brain* 103:221-244.

Rodriguez A, Whitson J, Granger R (2004) Derivation & analysis of basic computational operations of thalamocortical circuits. *J. Cog Neurosci*, 16: 856-877.

Rumelhart D, Zipser, D (1985) Feature discovery by competitive learning. *Cognitive Sci* 9:75-112.

Shimono K, Brucher F, Granger R, Lynch G, Taketani M (2000) Origins and distribution of cholinergically induced beta rhythms in hippocampal slices. *J Neurosci* 20:8462-8473.

Steriade M, Llinas R (1988) The functional states of thalamus and associated neuronal interplay. *Phys Rev* 68:649-742.

Szentagothai J (1975) The 'module-concept' in cerebral cortex architecture. *Brain Res* 95:475-496.

Valverde F (2002) Structure of cerebral cortex. Intrinsic organization & comparative analysis. *Rev Neurol* 34:758-780.

von der Malsburg C (1973) Self-organization of orientation sensitive cells in striate cortex. *Kybernetik* 14:85-100.

White E, Peters A (1993) Cortical modules in the posteromedial barrel subfield (Sml) of the mouse. *J Comp Neurol* 334:86-96.

Zhang S, Huguenard J, Prince D (1997) GABA_A receptor mediated Cl⁻ currents in rat thalamic reticular & relay neurons. *J Neurophysiol* 78:2280-2286.